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Arjona Antolín et al.

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# (54) CATALYST FOR OBTAINING HIGHER ALCOHOLS

(71) Applicant: **Abengoa Bioenergía Nuevas Technolgías, S.A.**, Seville (ES)

(72) Inventors: Ricardo Arjona Antolín, Seville (ES);

Juan Luís Sanz Yagüe, Seville (ES); Avelino Corma Canós, Valencia (ES); Marcelo Eduardo Domine, Valencia

(ES)

(73) Assignee: ABENGOA BIOENERGÍA NUEVAS

TECNOLOGÍAS, S.A., Seville (ES)

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Primary Examiner — Brian J Davis (74) Attorney, Agent, or Firm — Novak Druce Connolly Bove + Quigg LLP

## (57) ABSTRACT

The present invention relates to a process for obtaining metal oxide catalysts comprising gallium which are capable of synthesizing higher alcohols from lower alcohols. The process for obtaining said catalysts is also disclosed.

## 11 Claims, 2 Drawing Sheets

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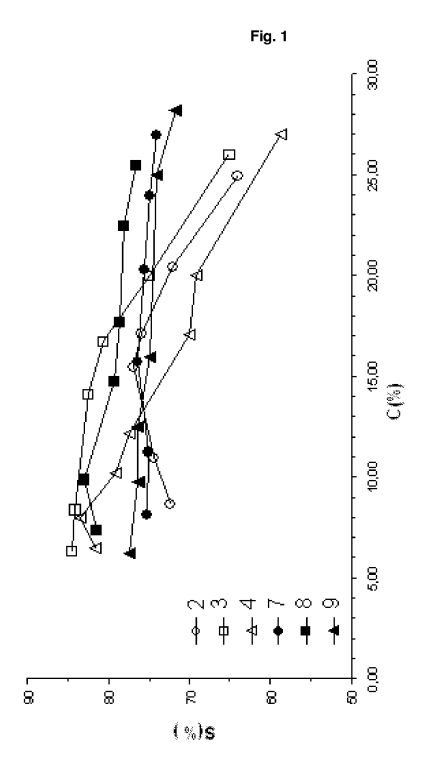
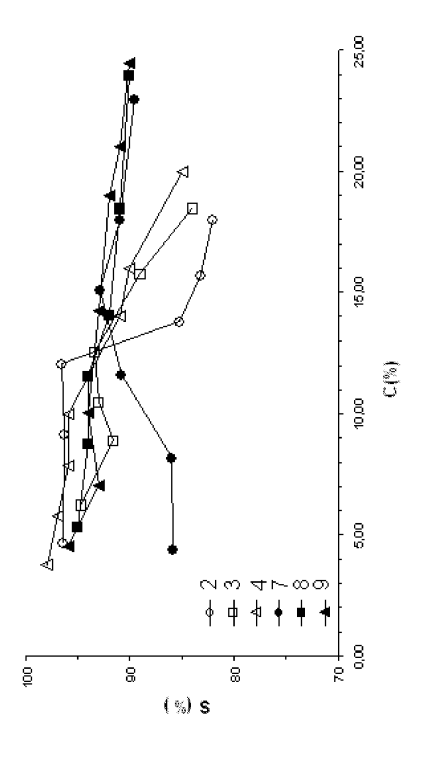


Fig. 2



## CATALYST FOR OBTAINING HIGHER ALCOHOLS

The present invention relates to hydrotalcite-derived catalysts that comprise gallium and a noble metal, as well as the 5 process for obtaining them and the use thereof in reactions designed to obtain higher alcohols. Therefore, the present invention belongs to the field of catalysts designed to obtain higher alcohols.

## BACKGROUND OF THE INVENTION

The synthesis of high-molecular-weight alcohols (containing between 4 and 8 carbon atoms) from low-molecularweight alcohols (e.g. methanol and ethanol) has acquired 15 considerable interest in recent years due to the potential use of these oxygenated compounds in the preparation of automotive fuels and also additives for the latter.

In particular, n-butanol (n-ButOH), with a worldwide production of approx. 4-5 million tons per year, is primarily used 20 catalysts are still needed for the synthesis thereof. as an additive for liquid fuels, as a starting chemical reagent in the plastics industry and also as an extraction agent in the aroma and fragrance industry. N-butanol is primarily produced through the petrochemical pathway. Methods for producing n-butanol by fermentation are also widely known.

Alternatively, n-butanol can be obtained by means of the well-known Guerbet reaction, which makes it possible to convert a low-molecular-weight alcohol (e.g. ethanol) into a linear or branched alcohol with a higher molecular weight in the presence of a metal alkoxide dissolved in the alcohol to be 30 transformed. The main disadvantages associated with the Guerbet reaction are: i) the production of water, which must be eliminated from the reaction medium in order to favour the formation of the desired compounds, ii) the production of carboxylic acids, and iii) the use of homogeneous catalysts, 35 which cause both corrosion in the reactors and final separation problems.

U.S. Pat. No. 5,300,695 discloses K, Na, Ba and Cs cations exchanged zeolites, amongst others, as catalysts in the condensation of low-molecular-weight alcohols, to obtain selec- 40 tivities to iso-butanol of 30%-35% at a 45% methanol conversion rate. Several basic oxides containing Cu, commonly used in the production of high-molecular-weight alcohols from syn-gas (CO/H<sub>2</sub>), have also been assayed in methanol and ethanol condensation reactions, to produce high-molecu- 45 lar-weight alcohols, albeit with a relatively low production of C<sub>4</sub> alcohols (U.S. Pat. No. 5,387,570). Another group of catalysts used are calcium-phosphate-based materials of the hydroxyapatite type (US20070255079). These catalysts have been tested in fixed-bed continuous reactors at 300° C. with 50 very short contact times. The best results were obtained for materials with a Ca/P molar ratio of 1.5-1.7, with ethanol conversion of 12% and selectivities to C<sub>4</sub> alcohols (primarily iso-butanol) close to 78%. Oxides of alkaline-earth metals, particularly materials based on MgO with supported transi- 55 tion metals, have also been used for the transformation of ethanol into n-butanol.

In recent years, numerous scientific publications and patents have appeared on the use of hydrotalcite-type materials as catalysts in alcohol condensation reactions, such as the 60 Guerbet reaction, in both batch systems and fixed-bed continuous reactors. The studies performed with these mixed Mg—Al oxides revealed that the catalytic activity of these materials is dependent on the nature, density and strength of the basic surface sites, which, in turn, are dependent on the 65 molar Mg/AI composition in the solid. For example, international application WO2009026510 discloses a process for

2

synthesising n-butanol by means of a material derived from the thermal decomposition of a hydrotalcite which preferably comprises magnesium and aluminum. Moreover, WO2009097312, US20100160693 and WO2009097310 disclose materials obtained by the thermal decomposition of hydrotalcites modified by the inclusion of metal carbonates and ethylenediamine-tetraacetates, which have been developed by DU PONT as catalysts in alcohol condensation reactions operating in a fixed bed at 300° C. and atmospheric pressure. The best results under these conditions have been achieved with a material derived from Mg—Al-based hydrotalcite (containing OH- as the anion), which presents ethanol conversion (≈44%) with moderate selectivities (≈44%) to n-butanol. When these same materials were assayed in the catalytic conversion of ethanol into n-butanol in the presence of hydrogen in the reaction system, the yields of n-butanol obtained were significantly lower in all cases.

Given the importance of higher alcohols, new, improved

## DESCRIPTION OF THE INVENTION

The present invention relates to a new metal oxide catalyst that comprises gallium and which is capable of obtaining higher alcohols from lower alcohols.

The invention presents the following advantages with respect to the catalysts in the state of the art:

at a given concentration of palladium, the hydrotalcitederived catalysts that comprise gallium in their structure provide higher yields of n-butanol than their analogues without gallium,

moreover, they also show a greater catalytic activity (measured through the TON, Turnover Number, parameter) than catalysts that present the same concentration of Pd but do not have gallium in their structure, which means that the catalysts are more stable under the reaction conditions, performing a greater number of catalytic cycles in a given reaction time,

a lower temperature is required in order to perform the process.

Therefore, one aspect of the present invention relates to a process for obtaining a catalyst (hereinafter process of the invention), which comprises the following steps:

a) thermal decomposition of a hydrotalcite with formula  $[M1_{1-(x+y)}M2_xM3_y (OH)_2][A^{m-}_{(x+y)/m}.nH_2O]$  to obtain a metal oxide, wherein:

M1 is at least one bivalent metal selected from the list that comprises Mg, Zn, Cu, Co, Mn, Fe, Ni and Ca,

M2 is at least one trivalent metal selected from the list that comprises Al, La, Fe, Cr, Mn, Co and Ni, M3 is trivalent Ga,

A is at least one anion selected from the list that comprises hydroxide, chloride, fluoride, bromide, iodide, nitrate, perchlorate, chlorate, bicarbonate, acetate, benzoate, methanesulfonate, p-toluenesulfonate, phenoxide, alkoxide, carbonate, sulfate, terephthalate, phosphate, hexacyanoferrate (III) and hexacyanoferrate (II),

x is a value between 0 and 0.5, preferably, x is a value between 0.1 and 0.5, and, even more preferably, between 0.1 and 0.4; y is a value between 0.00001 and 0.49, preferably between 0.00005 and 0.45, and, even more preferably, between 0.0001 and 0.4; m is an integer between 1 and 4; and n is greater than 0, preferably n is a value between 0 and 100, and, even more preferably, between 0 and 20. n indicates the number of crystallisation water molecules and is dependent on the composition of the hydrotalcite cations.

b) addition of at least one noble metal selected from the list that comprises Pd, Ru, Rh and Re to the solid obtained in step

"Bivalent metal" or "trivalent metal" is understood to mean a metallic cation with a +2 or +3 charge, respectively.

"Hydrotalcite" is understood to mean the structural family of laminar mixed hydroxides with the formula described above. The general structure of hydrotalcites is well-known to persons skilled in the art.

The term "thermal decomposition" is understood to mean 10 a chemical decomposition or structural change caused by the action of heat. This decomposition may be total or partial, depending on whether said decomposition is performed to completion or, on the contrary, is partially performed. This thermal decomposition may be performed at temperatures 15 greater than 150° C. and in the presence of an oxidising or a non-oxidising gas.

In a preferred embodiment of the process of the invention, there is a step (a') prior to (a), where the hydrotalcite is synthesised by the co-precipitation of at least one M1 com- 20 pound and at least one compound of a trivalent metal selected from the list that comprises M2 and M3; preferably, the hydrotalcite is obtained by the co-precipitation of M1, M2 and M3 compounds. The best results have been obtained when M3 is incorporated into the catalyst by co-precipitation 25 simultaneously with M1 and M2. Preferably, the co-precipitation is performed in the aqueous phase. The co-precipitation is preferably performed following the addition of a solution of at least one anion A selected from the list that comprises hydroxide, chloride, fluoride, bromide, iodide, nitrate, perchlorate, chlorate, bicarbonate, acetate, benzoate, methanesulfonate, p-toluenesulfonate, phenoxide, alkoxide, carbonate, sulfate, terephthalate, phosphate, hexacyanoferrate (III) and hexacyanoferrate (II) to a solution of at least one M1 compound and at least one compound of a trivalent metal 35 selected from the list that comprises M2 and M3, preferably a solution of M1, M2 and M3 compounds. This anion A may be introduced between the sheets of the resulting hydrotalcite. In order to obtain solutions of anion A, sodium and/or potassium salts thereof may be used. Preferably, A is at least one anion 40 event that M3 has not been incorporated into the hydrotalcite selected from the list that comprises carbonate, bicarbonate and hydroxide. The best results are obtained when the coprecipitation is performed at a pH higher than 7, preferably between 10 and 14. In order to regulate the pH, sodium and/or potassium hydroxide are preferably used.

Preferably, prior to the precipitation of said compounds, there is a dissolution of at least one M1 compound and at least one compound of a trivalent metal selected from the list that comprises M2 and M3, preferably a solution of M1, M2 and M3 compounds. Soluble M1, M2 and M3 compound is under- 50 stood to mean any salt which, when in contact with a solvent, preferably water, is dissociated. Examples of soluble M1, M2 and M3 compounds may be nitrates, halides, sulfates, carboxylates and, in general, oxoacids that comprise M1, M2 or

In a preferred embodiment of the process of the invention, M1 is Mg. Moreover, M2 is preferably Al. As regards the anion, A is preferably at least one anion selected from the list that comprises CO<sub>3</sub><sup>2-</sup>, HCO<sub>3</sub><sup>-</sup>, O<sub>2</sub><sup>-</sup>, OH<sup>-</sup>, Cl<sup>-</sup>, NO<sub>3</sub><sup>2-</sup>, Cl<sup>-</sup>, F<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>, ClO<sub>4</sub><sup>-</sup>, CH<sub>3</sub>COO<sup>-</sup>,  $C_6H_5COO^-$ , and SO<sub>4</sub><sup>2-</sup>; even 60 more preferably, CO<sub>3</sub><sup>2-</sup>, HCO<sub>3</sub><sup>-</sup>, O<sub>2</sub><sup>-</sup> and OH<sup>-</sup>.

In a preferred embodiment of the process of the invention, the thermal decomposition of hydrotalcite is performed by air calcination at a temperature ranging between 250° C. and 650° C., preferably between 350° C. and 550° C. The thermal decomposition of hydrotalcite is preferably performed for an interval ranging between 0.5 and 48 hours, preferably

between 1 and 24 hours, and, more preferably, between 1 and 6 hours. This process may be performed by heating the hydrotalcite in a gaseous atmosphere and may be performed in a static oven or a calcination reactor with a controlled gas flow, the latter being the preferred system. The gas may be an oxidising gas or a non-oxidising gas. Examples of oxidising gases may include air and oxygen. Examples of non-oxidising gases may be inert gases, such as nitrogen, argon, helium and reducing gases, such as, for example, carbon dioxide, hydrogen and ammonia. Preferably, the calcination is performed in the presence of oxygen, nitrogen or mixtures thereof, and, even more preferably, in the presence of oxygen and nitrogen.

The gels resulting from the co-precipitation as described above are filtered, washed with water and adequately dried. The presence of a hydrotalcite-type structure may be corroborated by means of X-ray diffraction analysis (XRD), whilst the composition (quantity and type of constituent) of the hydrotalcite or the corresponding mixed oxide obtained by the thermal decomposition of the aforementioned hydrotalcite may be determined by means of inductively coupled plasma mass spectrometry (ICP-MS) and chemical analysis, amongst others.

In a preferred embodiment of the process of the invention, the addition of the noble metal of step (b) is performed by wet impregnation, incipient volume impregnation or depositionprecipitation, preferably by incipient volume impregnation. The incipient volume impregnation method, also called incipient wetness impregnation method, is based on the use of a minimum quantity of liquid for the impregnation, only that which is necessary to reach the maximum saturation of the corresponding solid.

Another embodiment of the present invention is the process as described above, where the concentration of the noble metal in the metal oxide ranges between 0.001% and 10% by weight with respect to the total metal oxide, preferably between 0.01% and 5%.

In a particular embodiment of the present invention, in the in the co-precipitation step, M3 may be incorporated into the metal oxide in a step subsequent to the synthesis by means of wet impregnation, incipient volume impregnation and/or deposition-precipitation. This incorporation may be performed prior to or simultaneously with the addition of at least one noble metal selected from the list that comprises Pd, Ru and Rh. preferably Pd.

At a given concentration of palladium, the hydrotalcitederived catalysts that comprise gallium in their structure provide higher yields of n-butanol in a nitrogen atmosphere than their analogues without gallium. Furthermore, they also show a higher TON than catalysts that present the same concentration of Pd but do not have gallium in their structure. This fact is an indication of the greater stability of the catalysts of the invention under the reaction conditions.

Moreover, as may be observed in FIGS. 1 and 2, the catalysts of the invention present higher selectivities to butanol at a given ethanol conversion than the catalysts in the state of the

In a preferred embodiment of the process of the invention, M1 is Mg. Moreover, M2 is preferably Al. As regards the anion, A is preferably at least one anion selected from the list that comprises  ${\rm CO_3}^2$ ,  ${\rm HCO_3}^-$ ,  ${\rm O_2}^-$ ,  ${\rm OH}^-$ ,  ${\rm CI}^-$ ,  ${\rm NO_3}^2$ ,  ${\rm Cr}$ ,  ${\rm F}^-$ ,  ${\rm Br}^-$ ,  ${\rm I}^-$ ,  ${\rm CIO_4}^-$ ,  ${\rm CH_3COO}^-$ ,  ${\rm C_6H_5COO}^-$ , and  ${\rm SO_4}^{2-}$ ; even more preferably,  $CO_3^{2-}$ ,  $HCO_3^-$ ,  $O_2^-$  and  $OH^-$ .

In a preferred embodiment of the process of the invention, the noble metal that is added in step (b) is Pd. The best yields

of n-butanol have been obtained when the hydrotalcites containing Ga are impregnated with Pd.

In a preferred embodiment, the process of the invention further comprises a step (c), subsequent to (b), where the product obtained in step (b) is calcined. Preferably, this calcination is calcination in an atmosphere of oxygen, nitrogen or any mixture thereof. The calcination is preferably performed at a temperature ranging between 250° C. and 650° C., and, even more preferably, between 350° C. and 550° C. This calcination is preferably performed for an interval ranging between 0.5 and 48 hours, preferably between 1 and 24 hours, and, even more preferably, between 1 and 6 hours. This process may be performed by heating the hydrotalcite in a gaseous atmosphere and may be performed in a static oven or 15 a calcination reactor with a controlled gas flow, the latter being the preferred system. The gas may be an oxidising gas or a non-oxidising gas. Examples of oxidising gases may include air and oxygen. Examples of non-oxidising gases may be inert gases, such as nitrogen, argon, helium and reduc- 20 ing gases, such as, for example, carbon dioxide, hydrogen and ammonia. Preferably, the calcination is performed in the presence of oxygen, nitrogen or mixtures thereof, and, even more preferably, in the presence of oxygen and nitrogen.

further comprises a reduction step (d), subsequent to (c). During the reduction, the noble metal, which acts as one of the main active sites in the process, is reduced. The reduction is preferably performed in an H<sub>2</sub> atmosphere at a temperature ranging between 200° C. and 500° C., even more preferably 30 between 250° C. and 450° C. This reduction is preferably performed for an interval ranging between 0.5 and 48 hours, more preferably between 1 and 24 hours, and, even more preferably, between 1 and 6 hours. The reduction takes place immediately prior to the contact step with the reagent.

Another aspect of the present invention relates to a catalyst obtained by means of the process as described above (hereinafter catalyst of the invention).

At a given concentration of palladium, the catalysts of the invention provide higher yields of n-butanol than their ana- 40 ing higher alcohols, contact between the reagent and the logues without gallium. Moreover, they show a higher TON (Turnover Number) than the catalysts that present the same concentration of Pd but do not have gallium in their structure, which means that the catalysts are more stable under the reaction conditions, performing a greater number of catalytic 45 cycles prior to the deactivation thereof. Furthermore, they require a lower temperature to perform the process for obtaining higher alcohols.

Another aspect of the present invention relates to the use of the catalyst of the invention to obtain  $C_3$ - $C_{15}$  higher alcohols, 50 preferably C<sub>3</sub>-C<sub>8</sub>.

The term "C<sub>3</sub>-C<sub>15</sub> higher alcohols" is understood to mean any linear or branched alkyl chain with at least one hydroxyl functional group which has between 3 and 15 carbon atoms. Likewise, the term "C<sub>3</sub>-C<sub>8</sub> higher alcohols" is understood to 55 mean any linear or branched alkyl chain with at least one hydroxyl functional group which has between 3 and 8 carbon atoms. Thus, the higher alcohol will preferably be a  $C_3$ ,  $C_4$ ,  $C_5$ ,  $C_6$ ,  $C_7$  or  $C_8$ . Non-limiting examples would be propanol, isopropanol, n-butanol, 2-butanol, 2-methyl-2-butanol, 60 3-methyl-1-butanol-1-pentanol, 2-pentanol, 3-pentanol, 2,2dimethyl-1-propanol, 3-methyl-2-butanol, 1,5-pentanediol, 2,4-pentanediol, 2,2-dimethyl-1,3-propanediol, 1,2-butanediol, 1,3-butanediol, 1,4-butanediol, 2,3-butanediol, 1-heptanol, 2-heptanol, 3-heptanol, 4-heptanol, 2-methyl-2-hex- 65 2,2-dimethyl-3-pentanol-1-octanol, 3-octanol, 4-octanol, 2-ethyl-1-hexanol, 3-ethyl-1-hexanol

6

and 2,2-dimethyl-3-hexanol. Obviously, when the reagent is a C<sub>3</sub>, the higher alcohol obtained will be at least a C<sub>4</sub>, preferably a  $C_4$ ,  $C_5$ ,  $C_6$ ,  $C_7$  or  $C_8$ .

In a preferred embodiment of the use of the catalyst of the invention, the higher alcohol is preferably a  $C_4$ , more preferably n-butanol.

The obtainment of these higher alcohols is preferably performed by means of a process that comprises a contact step between at least one reagent selected from the list that comprises methanol, ethanol (EtOH), propanol and isopropanol, and the catalyst of the invention. Obviously, when the reagent is a C<sub>3</sub>, the higher alcohol obtained will be at least a C<sub>4</sub>, preferably a C<sub>4</sub>, C<sub>5</sub>, C<sub>6</sub>, C<sub>7</sub> or C<sub>8</sub>. Preferably, the reagent is ethanol.

Amongst other by-products, in the process for obtaining higher alcohols, C<sub>3</sub>-C<sub>20</sub> higher primary alcohols, preferably C<sub>4</sub>-C<sub>12</sub>, are also obtained. Moreover, C<sub>3</sub>-C<sub>20</sub> higher secondary alcohols, preferably C3-C11, may also be obtained. The hydroxyl function of said higher secondary alcohols will be preferably located at the C2. C2-C6 aldehydes may also be obtained. In particular, the main by-products are preferably ethanal, 2-butanol, butanal, 1-hexanol, 2-hexanol, hexanal, 1-octanol, 2-octanol and octanal.

In a preferred embodiment of the process for obtaining In a preferred embodiment, the process of the invention 25 higher alcohols, the contact between the reagent and the catalyst is performed in a reactor selected from the list that comprises discontinuous reactor, continuous stirred-tank reactor, fixed-bed continuous reactor and fluidized-bed continuous reactor, preferably a continuous reactor.

> In the particular embodiment wherein the reactor is a discontinuous reactor, contact between the reagent and the catalyst is performed at a temperature ranging between 50° C. and 450° C., preferably between 100° C. and 300° C. In this process, the weight ratio between the reagent and the catalyst is preferably between 2 and 200, preferably between 5 and 100. Moreover, it is performed for a time interval ranging between 2 minutes and 200 hours, preferably between 1 hour and 100 hours.

> In another preferred embodiment of the process for obtaincatalyst is performed in an atmosphere of nitrogen, argon, hydrogen or any mixture thereof, preferably in a nitrogenhydrogen atmosphere. Usually, higher selectivities to n-butanol are obtained in the presence of hydrogen.

> In another preferred embodiment of the process for obtaining higher alcohols, contact between the reagent and the catalyst is performed at a pressure of up to 120 bars, preferably between 20 and 80 bars.

> Throughout the description and the claims, the word "comprises" and the variants thereof are not intended to exclude other technical characteristics, additives, components or steps. For persons skilled in the art, other objects, advantages and characteristics of the invention will arise partly from the description and partly from the practise of the invention. The following examples and figures are provided for illustrative purposes, and are not intended to limit the scope of the present invention.

## BRIEF DESCRIPTION OF THE FIGURES

FIG. 1. Shows a comparative graph of the selectivities of the Pd/HT-4 and Pd/Ga-HT-4 catalysts in an N<sub>2</sub> atmosphere as a function of the conversion. S(%): Selectivity to but anol in mole %; C(%): ethanol conversion in mole %; 2: 1% Pd/HT-4 (Ex. 2); 3: 0.5% Pd/HT-4 (Ex. 3); 4; 0.25% Pd/HT-4 (Ex. 4); 7: 1% Pd/0.29% Ga-HT-4 (Ex. 7); 8: 0.50% Pd/0.29% Ga-HT-4 (Ex. 8); 9: 0.24% Pd/0.29% Ga-HT-4 (Ex. 9).

FIG. 2. Shows a comparative graph of the selectivities of the Pd/HT-4 and Pd/Ga-HT-4 catalysts in an N<sub>2</sub>—H<sub>2</sub> atmosphere as a function of the conversion. Legend same as in FIG. 1.

#### **EXAMPLES**

Below we will illustrate the invention by means of assays performed by the inventors, which demonstrate the efficacy of the hydrotalcite-derived catalysts that comprise gallium in  $^{10}$ their structure in the obtainment of higher alcohols.

## Example 1

## Synthesis of the HT-4 Catalyst (Mg/AI Molar Ratio≈4)

It was prepared by means of a standard co-precipitation process using two solutions. The first solution contained 36.45 g of Mg(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O and 13.60 g of Al(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O, <sup>20</sup> dissolved in 67.79 g of Milli-Q water, with a molar concentration of Al+Mg of 1.5. The second solution contained 12.53 g of NaOH and 16.16 g of Na<sub>2</sub>CO<sub>3</sub> in 89.63 g of Milli-Q water, and was used to produce the adequate precipitation of at ≈13. Both solutions were added, at a total flow velocity of 30 ml/h for approx. 4 h, to a container under vigorous stirring at room temperature. The gel formed was aged at room temperature for 1-2 h; thereafter, it was filtered and washed with distilled water until the carbonate was not detected in the  $^{30}$ filtered liquid (at pH≈7). Subsequently, the solid was dried in an oven at 60° C. for 14-16 h, and a mixed oxide called HT-4 was obtained, with a Mg/AI molar ratio≈3.8 and a surface area (BET method) of 257 m<sup>2</sup>/g. The BET method refers to the Brunauer-Emmett-Teller isotherm method.

#### Example 2

## Synthesis of the 1% Pd/HT-4 Catalyst

It was prepared from the material prepared as described in Example 1, wherein the incorporation of Pd (1.0% by weight, theoretical) into the HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.030 g of  $Pd(NH_3)_4Cl_2.6H_2O$  dissolved in 2.000 g of 45 Milli-Q water, to impregnate 1.014 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The 50 resulting Pd/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈1.00% by weight of Pd.

## Example 3

## Synthesis of the 0.5% Pd/HT-4 Catalyst

It was prepared from the material prepared as described in Example 1, wherein the incorporation of Pd (0.5%) by weight, theoretical) into the HT-4 material was performed by means 60 of the incipient wetness impregnation method, using, in this case, 0.015 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 2.000 g of Milli-Q water, to impregnate 1.023 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, 65 and, subsequently, it was reduced at 350° C. in an H2 atmosphere for 3 h prior to the catalytic application thereof. The

8

resulting Pd/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈0.50% by weight of Pd.

## Example 4

## Synthesis of the 0.25% Pd/HT-4 Catalyst

It was prepared from the material prepared as described in Example 1, wherein the incorporation of Pd (0.3% by weight, theoretical) into the HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.008 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 2.000 g of Milli-Q water, to impregnate 1.023 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H2 atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈0.25% by weight of Pd.

## Example 5

## Synthesis of the 0.80% Pt/HT-4 Catalyst

It was prepared from the HT-4 material prepared as the Al and Mg species, and to set the pH of the total mixture 25 described in Example 1, wherein the incorporation of Pt (1.0% by weight, theoretical) into the HT-4 material was performed by means of the incipient wetness impregnation method, using 0.025 g of H<sub>2</sub>Cl<sub>6</sub>Pt.6H<sub>2</sub>O dissolved in 2.000 g of Milli-Q water, to impregnate 1.025 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pt/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈0.80% by weight of Pt.

#### Example 6

## Synthesis of the 0.29% Ga-HT-4 Catalyst

It was prepared by means of a standard co-precipitation process using two solutions. The first solution contained 29.89 g of Mg(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O, 10.90 g of Al(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O and 0.06 g of Ga(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O, dissolved in 55.18 g of Milli-Q water, with a molar concentration of (Al+Mg+Ga) of 1.5. The second solution contained 12.52 g of NaOH and 10.52 g of Na<sub>2</sub>CO<sub>3</sub> in 72.60 g of Milli-Q water, and was used to produce the adequate precipitation of the Mg, Al and Ga species, and to set the pH of the total mixture at ≈13. Both solutions were added, at a total flow velocity of 30 ml/h for approximately 4 h, to a container under vigorous stirring at room temperature. The gel formed was aged at room temperature for 1-2 h; thereafter, it was filtered and washed with distilled water until the carbonate was not detected in the filtered liquid (at  $pH \approx 7$ ). Subsequently, the solid was dried in an oven at 60° C. for 14-16 h. The hydrotalcite (Ga-HT-4) obtained was calcined in air at 450° C. for 3-4 h, to obtain a mixed oxide with a Mg/AI molar ratio≈3.8, a Ga content of 0.29% by weight (measured by chemical analysis and ICP-MS), and a surface area (BET method) of 262  $\text{m}^2/\text{g}$ .

#### Example 7

## Synthesis of the 1% Pd/0.29% Ga-HT-4 Catalyst

It was prepared from the material prepared as described in Example 6, wherein the incorporation of Pd (1.0% by weight,

9

theoretical) into the Ga-HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.030 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 1.700 g of Milli-Q water, to impregnate 1.100 g of 0.29% Ga-HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/0.29% Ga-HT-4 material, characterised by chemical analysis and ICP-MS, contained  $\approx 1.00\%$  by weight of Pd.

## Example 8

#### Synthesis of the 0.50% Pd/0.29% Ga-HT-4 Catalyst

It was prepared from the material prepared as described in Example 6, wherein the incorporation of Pd (0.5% by weight, theoretical) into the Ga-HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.016 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 1.800 g of Milli-Q water, to impregnate 1.129 g of 0.29% Ga-HT-4. Once impregnated, the solid obtained was dried in an oven at  $100^{\circ}$  C. for 14-16 h; thereafter, it was calcined in air at  $450^{\circ}$  25 C. for 3-4 h, and, subsequently, it was reduced at  $350^{\circ}$  C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/0.29% Ga-HT-4 material, characterised by chemical analysis and ICP-MS, contained  $\approx 0.50\%$  by weight of Pd.

#### Example 9

## Synthesis of the 0.24% Pd/0.29% Ga-HT-4 Catalyst

It was prepared from the material prepared as described in Example 6, wherein the incorporation of Pd (0.3% by weight, theoretical) into the Ga-HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.008 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 1.800 g of Milli-Q water, to impregnate 1.011 g of 0.29% Ga-HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/0.29% Ga-HT-4 material, characterised by chemical analysis and ICP-MS, contained  $\approx$ 0.24% by weight of Pd.

## Example 10

#### Synthesis of the 1.06Pt/0.29% Ga-HT-4 Catalyst

It was prepared from the material prepared as described in 55 Example 6, wherein the incorporation of Pt (1.0% by weight, theoretical) into the Ga-HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.045 g of  $\rm H_2Cl_6Pt.6H_2O$  dissolved in 1.700 g of Milli-Q water, to impregnate 1.003 g of Ga-HT-4. Once 60 impregnated, the solid obtained was dried in an oven at  $100^\circ$  C. for 14-16 h; thereafter, it was calcined in air at  $450^\circ$  C. for 3-4 h, and, subsequently, it was reduced at  $350^\circ$  C. in an  $\rm H_2$  atmosphere for 3 h prior to the catalytic application thereof. The resulting Pt/0.29% Ga-HT-4 material, characterised by 65 chemical analysis and ICP-MS, contained  $\approx 1.06\%$  by weight of Pt.

10

## Example 11

## Synthesis of the 0.74% Pd/0.48% Ga/HT-4 Catalyst

It was prepared from the HT-4 material prepared as described in Example 1, wherein the incorporation of Ga (0.7% by weight, theoretical) into the HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.026 g of Ga(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O dissolved in 1.920 g of Milli-Q water, to impregnate 1.000 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h. The incorporation of Pd (1.0% by weight, theoretical) into the solid obtained was performed by means of the incipient wetness impregnation method, using, in this case, 0.095 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 1.400 g of Milli-Q water, to impregnate 1.540 g of the solid obtained in the first impregnation. Once impregnated, the final solid was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/Ga/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈0.74% by weight of Pd and ≈0.48% by weight of Ga.

## Example 12

#### Synthesis of the 0.74% Pd/0.29% Ga/HT-4 Catalyst

It was prepared from the HT-4 material prepared as 30 described in Example 1, wherein the incorporation of Ga (0.4% by weight, theoretical) into the HT-4 material was performed by means of the incipient wetness impregnation method, using, in this case, 0.015 g of Ga(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O dissolved in 1.920 g of Milli-Q water, to impregnate 1.000 g of HT-4. Once impregnated, the solid obtained was dried in an oven at 100° C. for 14-16 h. The incorporation of Pd (1.0% by weight, theoretical) into the solid obtained was performed by means of the incipient wetness impregnation method, using, in this case, 0.095 g of Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>.6H<sub>2</sub>O dissolved in 1.500 g of Milli-Q water, to impregnate 1.540 g of the solid obtained in the first impregnation. Once impregnated, the final solid was dried in an oven at 100° C. for 14-16 h; thereafter, it was calcined in air at 450° C. for 3-4 h, and, subsequently, it was reduced at 350° C. in an H<sub>2</sub> atmosphere for 3 h prior to the catalytic application thereof. The resulting Pd/Ga/HT-4 material, characterised by chemical analysis and ICP-MS, contained ≈0.74% by weight of Pd and ≈0.29% by weight of Ga.

#### Example 13

## Synthesis of the 5.0% Cu-HT-4 Catalyst

This catalyst was synthesised to illustrate hydrotalcite-type catalysts containing Cu, such as those cited in application WO2009026523. Various catalysts were synthesised with different concentrations of Cu, and the catalyst that provided the best results, in terms of selectivity and conversion, was selected in order to be compared to the catalysts of the invention

It was prepared by means of a standard co-precipitation process using two solutions. The first solution contained 28.73 g of Mg(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O, 10.50 g of Al(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O and 1.20 g of Cu(NO<sub>3</sub>)<sub>2</sub>.3H<sub>2</sub>O, dissolved in 56.25 g of Milli-Q water, with a molar concentration of (Al+Mg+Cu) of 1.5. The second solution contained 12.72 g of NaOH and 10.25 g of Na<sub>2</sub>CO<sub>3</sub> in 73.71 g of Milli-Q water, and was used to produce the adequate precipitation of the Mg, Al and Cu species, and

to set the pH of the total mixture at  $\approx$ 13. Both solutions were added (total flow velocity=30 ml/h for approximately 4 h) to a container under vigorous stirring at room temperature. The gel formed was aged at room temperature for 1-2 h; thereafter, it was filtered and washed with distilled water until the carbonate was not detected in the filtered liquid (at pH $\approx$ 7). Subsequently, the solid was dried in an oven at 60° C. for 14-16 h. The hydrotalcite (Cu-HT-4) obtained was calcined in air at 450° C. for 3-4 h, to obtain a mixed oxide with a Mg/AI molar ratio $\approx$ 3.8, a Cu content of  $\approx$ 5.0% by weight, characterised by chemical analysis and ICP-MS.

#### Example 14

## Synthesis of the 3.5% Co-HT-4 Catalyst

This catalyst was synthesised to illustrate hydrotalcite-type catalysts containing Co, such as those cited in application US20100160693. Various catalysts were synthesised with different concentrations of Co, and the catalyst that provided the best results, in terms of selectivity and conversion, was selected in order to be compared to the catalysts of the invention.

It was prepared by means of a standard co-precipitation process using two solutions. The first solution contained 25  $28.82 \text{ g of Mg(NO}_3)_2.6H_2O$ ,  $14.05 \text{ g of Al(NO}_3)_3.9H_2O$  and 1.17 g of Co(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O dissolved in 58.54 g of Milli-Q water, with a molar concentration of (Al+Mg+Cu) of 1.5. The second solution contained 13.81 g of NaOH and 10.87 g of Na<sub>2</sub>CO<sub>3</sub> in 77.91 g of Milli-Q water, and was used to produce <sup>30</sup> the adequate precipitation of the Mg, Al and Cu species, and to set the pH of the total mixture at ≈13. Both solutions were added (total flow velocity=30 ml/h for approximately 4 h) to a container under vigorous stirring at room temperature. The gel formed was aged at room temperature for 1-2 h; thereafter, 35 it was filtered and washed with distilled water until the carbonate was not detected in the filtered liquid (at pH≈7). Subsequently, the solid was dried in an oven at 60° C. for 14-16 h. The hydrotalcite (Co-HT-4) obtained was calcined in air at 450° C. for 3-4 h, to obtain a mixed oxide with a Mg/AI molar 40 ratio≈3.8, a Co content of ≈3.5% by weight, characterised by chemical analysis and ICP-MS.

## Example 15

## Synthesis of the 2.5% Ni-HT-4 Catalyst

This catalyst was synthesised to illustrate hydrotalcite-type catalysts containing Ni, such as those cited in application US20100160693. Various catalysts were synthesised with 50 different concentrations of Ni, and the catalyst that provided the best results, in terms of selectivity and conversion, was selected in order to be compared to the catalysts of the invention.

It was prepared by means of a standard co-precipitation 55 process using two solutions. The first solution contained 29.71 g of Mg(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O, 10.81 g of Al(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O and 0.78 g of Ni(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O, dissolved in 56.54 g of Milli-Q water, with a molar concentration of (Al+Mg+Cu) of 1.5. The second solution contained 12.85 g of NaOH and 10.37 g of 60 Na<sub>2</sub>CO<sub>3</sub> in 74.33 g of Milli-Q water, and was used to produce the adequate precipitation of the Mg, Al and Ni species, and to set the pH of the total mixture at  $\approx$ 13. Both solutions were added (total flow velocity=30 ml/h for approximately 4 h) to a container under vigorous stirring at room temperature. The 65 gel formed was aged at room temperature for 1-2 h; thereafter, it was filtered and washed with distilled water until the car-

12

bonate was not detected in the filtered liquid (at pH $\approx$ 7). Subsequently, the solid was dried in an oven at 60° C. for 14-16 h. The hydrotalcite (Ni-HT-4) obtained was calcined in air at 450° C. for 3-4 h, to obtain a mixed oxide with a Mg/AI molar ratio  $\approx$ 3.8, a Co content of  $\approx$ 2.5% by weight, characterised by chemical analysis and ICP-MS.

#### Example 16

Comparative Catalytic Activity of the Catalysts of Examples 1, 2, 5, 7 and 13-15 Under N<sub>2</sub> Atmosphere

3500 mg of ethanol and 200 mg of one of the catalytic materials of Examples 1, 2, 5, 7 and 13-15 were introduced 15 into a 12-ml stainless steel autoclave reactor, with a Tefloncoated inside and a magnetic stirrer. The reactor was hermetically closed, and the system contained a connector to a pressure meter (manometer), another connector for the loading of gases and a third outlet which made it possible to take samples at different time intervals. The reactor was initially pressurised with 24 bars of N<sub>2</sub>, and heated to 200° C. under continuous stirring, until the total system pressure reached approx. 30 bars (reaction time=0). Liquid samples (≈50 µl) were taken at different time intervals until 17-24 hours of reaction. The samples were filtered and diluted in a 2% by weight of chlorobenzene in acetonitrile standard solution, and analysed by means of gas chromatography in a GC-3900 Varian equipped with an FID detector and a 60-m TRB-624 capillary column; the ethanol conversion, in molar percentage (EtOH conv.), was calculated from the composition of the mixture obtained:

(initial moles of reagent-final moles of reagent)/(initial moles of reagent\*100)

The selectivities to n-butanol obtained were also calculated in molar percentage (Select. n-ButOH):

(moles of n-butanol/moles of total products)\*100

The total yield of n-butanol (Yield n-ButOH) was calculated as:

(EtOH conv.\*Select. n-ButOH)/100

In this manner, the following results were obtained:

TABLE 1

Catalytic activity of different mixed metal oxides in the transformation of ethanol into n-butanol under nitrogen atmosphere.

Ex. Catalyst	T (h)	EtOH conv.	Select. n-ButOH	Yield n-ButOH	TON <sup>a</sup>
1 HT-4	5	3.6	50.5	1.82	1
2 1.00% Pd/HT-4	5	15.5	76.9	11.9	629
5 0.80% Pt/HT-4	5	13.0	82.0	10.7	1268
7 1% Pd/0.29% Ga-HT-4	5	15.8	76.5	12.1	630
13 5.0% Cu-HT-4	5	8.3	68.8	5.92	42
14 3.5% Co-HT-4	5	1.2	76.2	0.91	8
15 2.5% Ni-HT-4	5	2.0	65.2	1.304	19

<sup>a</sup>TON = Turn Over Number in (mol/mol Pd or Pt).

These results show that the catalyst impregnated with Pd or Pt give better yields than the catalyst comprising co, Ni or Cu.

## Example 17

Comparative Catalytic Activity of the Catalysts of Examples 2-5 and 7-12 Under N<sub>2</sub> Atmosphere

3500 mg of ethanol and 200 mg of one of the catalytic materials of Examples 2-5 to 7-12 were introduced into a

12-ml stainless steel autoclave reactor, with a Teflon-coated inside and a magnetic stirrer. The reactor was hermetically closed, and the system contained a connector to a pressure meter (manometer), another connector for the loading of gases and a third outlet which made it possible to take samples 5 at different time intervals. The reactor was initially pressurised with 24 bars of N<sub>2</sub>, and heated to 200° C. under continuous stirring, until the total system pressure reaches approx. 30 bars (reaction time=0). Liquid samples (≈50 ρμl) were taken at different time intervals until 17-24 hours of reaction. The samples were filtered and diluted in a 2% by weight of chlorobenzene in acetonitrile standard solution, and analysed by means of gas chromatography in a GC-3900 Varian equipped with an FID detector and a 60-m TRB-624 capillary column; the ethanol conversion, in molar percent- 15 age (EtOH conv.), was calculated from the composition of the mixture obtained:

(initial moles of reagent-final moles of reagent)/(initial moles of reagent\*100)

The selectivities to n-butanol obtained were also calculated in molar percentage (Select. n-ButOH):

(moles of n-butanol/moles of total products)\*100

The total yield of n-butanol (Yield n-ButOH) was calculated as:

(EtOH conv.\*Select. n-ButOH)/100

In this manner, the following results were obtained:

TABLE 2

Catalytic activity of different mixed metal oxides in the transformation

of ethanol into n-butanol under nitrogen atmosphere.

Ex. Catalyst	T (h)	EtOH conv.	Select. n-ButOH	Yield n- ButOH	$TON^a$	35
2 1% Pd/HT-4	5	15.5	76.9	11.9	629	
	17	17.2	75.8	13.0	695	
3 0.5% Pd/HT-4	5	14.1	82.5	11.7	1145	
	17	16.8	80.6	13.5	1359	
4 0.25% Pd/HT-4	5	10.2	79.0	8.1	2073	40
	17	12.2	77.5	9.4	2460	-1
5 0.80% Pt/HT-4	5	13.0	82.0	10.7	1268	
	17	14.2	50.5	7.2	1385	
7 1% Pd/0.29% Ga-HT-4	5	15.8	76.5	12.1	630	
	17	20.4	75.5	15.4	825	
8 0.5% Pd/0.29% Ga-HT-4	5	14.8	79.3	11.7	1195	4.
	17	17.7	78.6	13.9	1435	4:
9 0.24% Pd/0.29% Ga-HT-4	5	12.5	76.4	9.6	2539	
	17	16.0	74.8	12.0	3237	
10 1.06% Pt/0.29% Ga-HT-4	5	12.0	69.0	8.3	881	
	17	13.8	56.6	7.8	1013	
11 0.74% Pd/0.48% Ga/HT-4	5	14.5	65.0	9.4	829	
	17	18.1	62.1	11.2	1034	50

<sup>&</sup>lt;sup>a</sup>TON = Turn Over Number in (mol/mol Pd or Pt).

12 0.74% Pd/0.29% Ga/HT-4

These results show that, at a given concentration of palladium, the hydrotalcite-derived catalysts that comprise gallium in their structure provide higher yields of n-butanol under nitrogen atmosphere than their analogues without gallium. Moreover, they also show a higher TON than the catalysts possessing the same concentration of Pd but without gallium in their structure. This fact is an indication of the greater stability of the catalysts of the invention under the reaction conditions.

13.0

16.3

17

63.0

60.0

8.2

9.8

743

931

Moreover, as can be observed in FIG. 1, the catalysts of the invention show higher selectivities to but anol under  $N_2$  atmosphere at a given ethanol conversion value than the catalysts in the state of the art.

14

Another fact worth mentioning is that the catalysts of the invention make it possible to achieve lower concentrations of Pd whilst maintaining high yields of n-butanol, as compared to the catalysts in the state of the art.

#### Example 18

Comparative Catalytic Activity of the Catalysts of Examples 2-5 and 7-12 Under N<sub>2</sub>—H<sub>2</sub> Atmosphere

3500 mg of ethanol and 200 mg of one of the catalytic materials of Examples 2-5 and 7-12 were introduced into a 12-ml stainless steel autoclave reactor, with a Teflon-coated inside and a magnetic stirrer. The reactor was hermetically closed, and the system contained a connector to a pressure meter (manometer), another connector for the loading of gases and a third outlet which made it possible to take samples at different time intervals. The reactor was initially pressurised with 10 bars of hydrogen, and, thereafter, was taken to a total pressure of 24 bars by adding N<sub>2</sub>. Subsequently, it was heated to 200° C. under constant stirring, until the total system pressure reaches approx. 32-33 bars (reaction time=0). Liquid samples (≈50 μl) were taken at different time intervals until 17-24 hours of reaction. The samples were filtered and diluted in a 2% (W/W) of chlorobenzene in acetonitrile standard solution, and analysed by means of gas chromatography in a GC-3900 Varian equipped with an FID detector and a 60-m TRB-624 capillary column; the ethanol conversion, in molar percentage (EtOH conv.), was calculated from the composition of the mixture obtained:

(initial moles of reagent-final moles of reagent)/(initial moles of reagent\*100)

The selectivities to n-butanol obtained were also calculated in molar percentage (Select. n-ButOH):

(moles of n-butanol/moles of total products)\*100

The total yield of n-butanol (Yield n-ButOH) was calculated as:

(EtOH conv.\*Select. n-ButOH)/100

10 1.06% Pt/0.29% Ga-HT-4

In this manner, the following results were obtained:

TABLE 3

Catalytic activity of different mixed metal oxides in the transformation

of ethanol into n-butanol under nitrogen and hydrogen atmosphere.

Ex. Catalyst	T (h)	EtOH conv.	Select. n-ButOH	Yield n- ButOH	$TON^{\alpha}$
2 1% Pd/HT-4	5	12.1	95.8	11.6	488
	17	13.8	83.2	11.5	560
3 0.5% Pd/HT-4	5	10.5	93.0	9.8	849
	17	12.5	93.3	11.7	1015
4 0.25% Pd/HT-4	5	7.8	96.0	7.5	1585
	17	10.0	96.0	9.6	2025
5 0.80% Pt/HT-4	5	5.0	87.0	4.4	488
	17	8.0	85.0	6.8	780
7 1% Pd/0.29% Ga-HT-4	5	11.7	90.9	10.6	472
	17	15.1	92.8	14.0	612
8 0.5% Pd/0.29% Ga-HT-4	5	11.6	94.0	10.9	937
	17	14.1	92.0	13.0	1141
9 0.24% Pd/0.29% Ga-HT-4	5	10.1	94.0	9.4	2034
	17	14.3	93.0	13.3	2888

96.0

7.7

9.9

587

763

Catalytic activity of different mixed metal oxides in the transformation of ethanol into n-butanol under nitrogen and hydrogen atmosphere.

Ex. Catalyst	T (h)	EtOH conv.	Select. n-ButOH	Yield n- ButOH	$TON^a$	5
11 0.74% Pd/0.48% Ga/HT-4	5	7.0	85.0	6.0	400	
	17	9.7	85.1	8.3	554	
12 0.74% Pd/0.29% Ga/HT-4	5	6.0	87.0	5.2	343	
	17	10.8	87.6	9.5	617	10

<sup>&</sup>quot;TON = Turn Over Number in (mol/mol Pd or Pt).

These results show that, at a given concentration of palladium, the hydrotalcite-derived catalysts that comprise gallium in their structure provide higher yields of n-butanol 15 under nitrogen and hydrogen atmosphere than their analogues without gallium. Moreover, they also show a higher TON than the catalysts possession the same concentration of Pd but without gallium in their structure. This fact is an indication of the greater stability of the catalysts of the invention under the reaction conditions. On the other hand, it is worth noting that incorporating hydrogen into the reaction mixture increases the selectivities to butanol.

Moreover, as can be observed in FIG. 2, the catalysts of the invention show higher selectivities to butanol in an  $N_2$ — $H_2$  25 atmosphere at a given ethanol conversion value than the catalysts in the state of the art.

Another fact worth mentioning is that the catalysts of the invention make it possible to achieve lower concentrations of Pd whilst maintaining high yields of n-butanol, as compared 30 to the catalysts in the state of the art.

The invention claimed is:

- 1. A process for obtaining a catalyst, which comprises the following steps:
  - a) thermal decomposition of a hydrotalcite with the formula  $[M1_{1-x(x+y)}M2_xM3_y(OH)_2][A^{m-}_{(x+y)/m}.nH_2O]$ , to obtain a metal oxide, wherein:
    - M1 is at least one bivalent metal selected from the group consisting of Mg, Zn, Cu, Co, Mn, Fe, Ni and Ca,
    - M2 is at least one trivalent metal selected from the group consisting of Al, La, Fe, Cr, Mn, Co and Ni,

M3 is trivalent Ga,

16

- A is at least one anion selected from the group consisting of hydroxide, chloride, fluoride, bromide, iodide, nitrate, perchlorate, chlorate, bicarbonate, acetate, benzoate, methanesulfonate, p-toluenesulfonate, phenoxide, alkoxide, carbonate, sulfate, terephthalate, phosphate, hexacyanoferrate (III) and hexacyanoferrate (II),
- x is a value between 0 and 0.5:
- y is a value between 0.00001 and 0.49;
- m is an integer between 1 and 4; and
- n is greater than 0, and
- b) addition of Pd to the solid obtained in step (a);
  - wherein the hydrotalcite is obtained by the co-precipitation of M1, M2 and M3 compounds.
- 2. The process according to claim 1, wherein the co-precipitation is performed by the addition of a solution of at least one compound selected from the group consisting of carbonate, bicarbonate and hydroxide to a solution of M1, M2 and M3 compounds.
- 3. The process according to claim 1, wherein the thermal decomposition of step (a) is calcination in an atmosphere of oxygen, nitrogen or any mixture thereof at a temperature ranging between 250° C. and 650° C.
- **4**. The process according to claim **1**, wherein the addition of Pd of step (b) is performed by wet impregnation, incipient volume impregnation or deposition-precipitation.
  - 5. The process according to claim 1, wherein M1 is Mg.
  - 6. The process according to claim 1, wherein M2 is Al.
- 7. The process according to claim 1, where A is at least one anion selected from the group consisting of list that comprises  ${\rm CO_3}^2$ ,  ${\rm HCO_3}^-$ ,  ${\rm O_2}^-$  and  ${\rm OH}^-$ .
- 8. The process according to claim 1, which further comprises a step (c), subsequent to (b), where the product obtained in step (b) is calcined.
- 9. The process according to claim 8, which further comprises a reduction step (d), subsequent to (c).
- 10. A catalyst obtained by means of the process according to claim 1
- 11. A process to obtain n butanol comprising a contact step between ethanol and the catalyst described according to claim

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